## IN THE CLAIMS

The claims are as follows:

- 1. (Original) A non-human animal having a neurologic disease induced by the process of: perfusing the non-human animal with a pharmacologically effective amount of a combination of an Aβ compound, at least one pro-oxidative compound, and at least one anti-oxidant inhibitor, wherein the perfusion produces impaired performance of the animal in memory and learning tests and induces abnormal neuropathology in a brain of the animal, wherein said impaired performance and abnormal neuropathology are in comparison with control non-human animals.
- 2. (Original) The non-human animal of claim 1, wherein the A $\beta$  compound comprises A $\beta_{42}$ .
- 3. (Original) The non-human animal of claim 1, wherein the A $\beta$  compound comprises a peptide fragment of A $\beta$ <sub>42</sub>.
- 4. (Original) The non-human animal of claim 3, wherein the peptide fragment of  $A\beta_{42}$  comprises at least one of  $A\beta_{1-40}$  or  $A\beta_{24-35}$ .
- 5. (Original) The non-human animal of claim 1, wherein the A $\beta$  compound comprises a peptidomimetic that mimicks A $\beta$ <sub>42</sub>.
- 6. (Original) The non-human animal of claim 1, wherein the at least one prooxidative compound is selected from the group consisting of ferrous sulfate, copper sulfate, cobalt sulfate, manganese sulfate, and zinc sulfate.
  - 7. (Original) The non-human animal of claim 1, wherein the at least one pro-

Title: ANIMAL MODEL SIMULATING NEUROLOGIC DISEASE

oxidative compound comprises ferrous sulfate.

(Original) The non-human animal of claim 1, wherein the at least one anti-oxidant 8. inhibitor comprises buthionine sulfoximine.

Page 3

Dkt: 1941.012US1

- (Original) The non-human animal of claim 1, wherein the process further 9. comprises perfusing the non-human animal with an effective amount of a phosphatase inhibitor.
- (Original) The non-human animal of claim 9, wherein the phosphatase inhibitor is 10. selected from the group consisting of okadaic acid, 1-nor-okadaone, bioallethrin, calycullin A, cantharidic acid, cantharidin, cypermethrin, deltamethrin, endothall, endothall thioanhydride, fenvalerate, okadol, permethrin, phenylarsine oxide, pyrophosphate, sodium fluoride, and vanadate.
- (Original) The non-human animal of claim 9, wherein the phophatase inhibitor 11. comprises okadaic acid.
- (Original) The non-human animal of claim 1, wherein the process further 12. comprises perfusing the non-human animal with an effective amount of a pro-inflammatory compound.
- (Original) The non-human animal of claim 12, wherein the pro-inflammatory 13. compound is selected from the group consisting of TNF-α, IL-6, and IL-1b.
- (Original) The non-human animal of claim 12, wherein the pro-inflammatory 14. compound comprises TNF- $\alpha$ .
- 15. (Original) A method for inducing a neurologic disease in a non-human animal, comprising:

perfusing the non-human animal with a pharmacologically effective amount of a

Page 4 Dkt: 1941.012US1

combination of an AB compound, at least one pro-oxidative compound, and at least one antioxidant inhibitor.

- (Original) The method of claim 15, wherein the A $\beta$  compound comprises A $\beta_{42}$ . 16.
- (Original) The method of claim 15, wherein the AB compound comprises a 17. peptide fragment of  $A\beta_{42}$ .
- 18. (Original) The method of claim 17, wherein the peptide fragment of  $A\beta_{42}$ comprises at least one of  $A\beta_{1-40}$  or  $A\beta_{24-35}$ .
- (Original) The method of claim 15, wherein the A\beta compound comprises a 19. peptidomimetic that mimicks Aβ<sub>42</sub>.
- (Original) The method of claim 15, wherein the at least one pro-oxidative 20. compound is selected from the group consisting of ferrous sulfate, copper sulfate, cobalt sulfate, manganese sulfate, and zinc sulfate.
- (Original) The method of claim 15, wherein the at least one pro-oxidative 21. compound comprises ferrous sulfate.
- (Original) The method of claim 15, wherein the at least one anti-oxidant inhibitor 22. comprises buthionine sulfoximine.
- (Original) The method of Claim 15, further comprising perfusing the non-human 23. animal with an effective amount of a phosphatase inhibitor.
- 24. (Original) The method of claim 23, wherein the phosphatase inhibitor is selected from the group consisting of okadaic acid, 1-nor-okadaone, bioallethrin, calycullin A, cantharidic

Dkt: 1941.012US1

acid, cantharidin, cypermethrin, deltamethrin, endothall, endothall thioanhydride, fenvalerte, okadol, perrnethrin, phenylarsine oxide, pyrophosphate, sodium fluoride, and vanadate.

- (Original) The method of claim 23, wherein the phophatase inhibitor comprises 25. okadaic acid.
- (Original) The method of claim 15, further comprising perfusing the non-human 26. animal with an effective amount of a pro-inflammatory compound.
- 27. (Original) The method of claim 27, wherein the pro-inflammatory compound is selected from the group consisting of TNF- $\alpha$ , IL-6, and IL-1b.
- (Original) The method of claim 27, wherein the pro-inflammatory compound 28. comprises TNF- $\alpha$ .
- (Original) A method of screening for an agent that ameliorates symptoms of a 29. neurologic disease, said method comprising:

comparing performance on memory and learning tests of a first non-human animal contacted with the agent with that of a second non-human animal not contacted with the agent, wherein the first and said second non-human animals have been co-infused with a pharmacologically effective amount of AB, at least one pro-oxidative compound, and at least one anti-oxidant inhibitor wherein the co-infusion produces impaired performance on the memory and learning tests and abnormal neuropathology in a brain of the first and second non-human animals, wherein the impaired performance and the abnormal neuropathology are in comparison with control non-human animals, whereby an agent which ameliorates the symptoms is identified by superior performance of said first non-human animal in comparison with the second nonhuman animal on the memory and learning tests.

(Original) A method for screening for an agent useful for treating a neurologic 30. disease, said method comprising:

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Title: ANIMAL MODEL SIMULATING NEUROLOGIC DISEASE

Page 6 Dkt: 1941.012US1

comparing performance on memory and learning tests of a first non-human animal contacted with the agent with that of a second non-human animal not contacted with the agent, wherein the first and said second non-human animals have been co-infused with a pharmacologically effective amount of Aβ and at least one pro-oxidative compound, and at least one anti-oxidant inhibitor, wherein the co-infusion produces impaired performance on the memory and learning tests and abnormal neuropathology in a brain of the first and second non-human animals, wherein the impaired performance and the abnormal neuropathology are compared with control non-human animals; and comparing neuropathology in the brain of the first and the second non-human animal when said first non-human animal exhibits superior performance on the memory and learning tests compared with the second non-human animal, whereby an agent which is useful for treating a neurologic disease is identified by a decrease in neuropathologic findings in the first non-human animal in comparison with the second non-human animal.